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Stapedial Artery: From Embryology to Different Possible Adult Configurations

 S. Bonasia,  S. Smajda,  G. Ciccio, and  T. Robert



ABSTRACT

SUMMARY: The stapedial artery is an embryonic artery that represents the precursor of some orbital, dural, and maxillary branches. Although its embryologic development and transformations are very complex, it is mandatory to understand the numerous anatomic variations of the middle meningeal artery. Thus, in the first part of this review, we describe in detail the hyostapedial system development with its variants, referring also to some critical points of ICA, ophthalmic artery, trigeminal artery, and inferolateral trunk embryology. This basis will allow the understanding of the anatomic variants of the middle meningeal artery, which we address in the second part of the review.

ABBREVIATIONS: MMA = middle meningeal artery; OA = ophthalmic artery; SA = stapedial artery

The stapedial artery (SA) is an embryologic artery that allows the development of orbital and dural arteries, and also of the maxillary branches. Its complex embryologic development explains numerous anatomic variations of the middle meningeal artery (MMA) and orbital arteries. Few anatomists¹⁻⁷ have dissected a human middle ear that bore a persistent SA and described the origin and course of this artery. Ear, nose, and throat surgeons reported cases discovered during middle ear surgeries, which explains the technical difficulties of the surgery in the presence of a persistent SA.⁸⁻²⁰ More recently, neuroradiologists have paid particular attention to this anatomic variant.²¹⁻²⁶ These studies, associated with the comparative anatomy,²⁷⁻²⁹ allow the understanding of the persistence of the SA. In this article, after a quick refresh concerning hyostapedial, carotid, and ophthalmic systems, the anatomic variations, including the SA, are presented and explained.

History

The first cadaveric case of persistent SA was described by Hyrtl³⁰ in 1836, who called attention to an artery that runs across the

obturator of the stapes in a human cadaver, with some similarities with a vessel found in hibernating animals. In the first half of the 20th century, the phenomenal publication by Padget,³¹ based on the dissections of 22 human embryos of the Carnegie collection, provided a great deal of information about the embryologic development of the craniofacial arteries and, in particular, of the hyostapedial system. In the same period, Altmann³² furnished a comprehensive explanation of the development of the aortic arches and of the carotid system variants. However, not until the 1960s, do we find the first angiographic demonstration of MMA variations and, in particular, the persistence of an SA.^{33,34} At the end of the 1970s, Lasjaunias was able to give a comprehensive explanation of all variations that implicate the SA.^{23,24} This group's more famous articles^{23-25,35-38} were published between 1975 and 1990 and are summarized in their textbook.³⁹ Diamond^{5,27,28} also published some articles based on the comparative anatomy, principally with the great apes, that increased our understanding of the stapedial system. In these past 3 decades, a few case reports of interesting anatomic variations that implicate the SA were published.^{17,40-47}

Comparative Anatomy

The studies of comparative anatomy were fundamental in the understanding of the embryologic development of the SA. At the beginning of the 20th century, Tandler,⁴⁸ when studying the SA development in rats, explained the SA regression and the role of the external carotid artery in the formation of the MMA. Rats differed from human embryos because the supraorbital branch of the SA is primarily dependent on the first aortic arch (mandibular branch). Fuchs,⁴⁹ in the rabbit, also showed that the first 2

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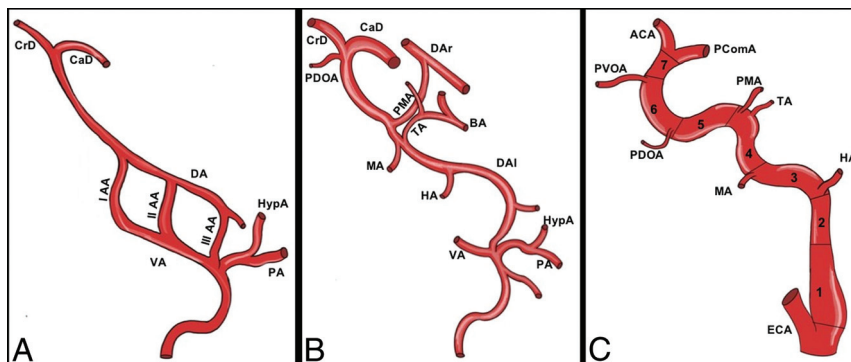


FIG 1. Embryologic segments of the ICA. Consecutive stages of ICA embryologic development. The first stages of development (A) are characterized by the presence of 3 aortic arches that link the ventral aorta (VA) and the dorsal aorta (DA). The VA regresses together with the ventral part of the aortic arches. The dorsal remnants of the aortic arches persist as embryonic arteries. These embryonic arteries divide the ICA into 7 embryologic segments: 1) the cervical segment: it derives from the remnant of the third aortic arch (III AA); 2) the ascending intrapetrous segment: it is the remnant of the DA between the second (II AA) and third (III AA) aortic arches; the division point between segments 2 and 3 is at the point of origin of the hyoid artery (HA), that is, the dorsal remnant of the second aortic arch (II AA); 3) the horizontal intrapetrous segment: it is the remnant of the DA between the first (I AA) and second (II AA) aortic arches; the division point is at the point of origin of the mandibular artery (MA), which corresponds to the dorsal remnant of the first aortic arch (I AA); 4) the intracavernous ascending segment: it originates from the DA between the first aortic arch (I AA) and the primitive maxillary artery (PMA), which connects the DA of the 2 sides (dorsal aorta left [DAL]; dorsal aorta right [DAR]). The trigeminal artery originates at the junction between the segment 4 and 5 of the ICA; this latter represents a primitive connection between the cavernous ICA and the basilar artery (BA); 5) the horizontal intracavernous segment: it derives from the DA between the PMA and the primitive dorsal OA (PDOA); 6) the clinoid segment: it corresponds to the DA between the PDOA and the primitive ventral OA; and 7) the terminal segment: the terminal ICA between primitive ventral OA and the primitive ICA bifurcation into the future anterior cerebral artery (ACA) and the future posterior communicating artery (PComA). The figure also shows the hypoglossal artery (HypA) and the proatlantal artery (PA), which originate proximal to the third aortic arch and will contribute to the formation of the external carotid artery (ECA) branches.

aortic arches participate in the formation of the external carotid artery as in human embryos. More recently, Diamond²⁸ and Falk²⁹ compared the development of the SA between human and great ape embryos and highlighted some analogies.

Embryology

The development and regression of the SA is strictly related to some embryonic systems. In this part of our article, we summarize the main steps of the ICA and ophthalmic artery (OA) development. We present a more-detailed description of the hyostapedial system to understand the different possible adult variations related to this artery.

The Carotid System. Because a detailed description of the ICA embryologic development is beyond the scope of this article, we focused only on the understanding of the “aberrant flow of the ICA.” It consists of an intratympanic course of the ICA and could be associated with a persistent SA and consequent variation in the origin of the MMA. The principal steps of ICA development are summarized in Fig 1.

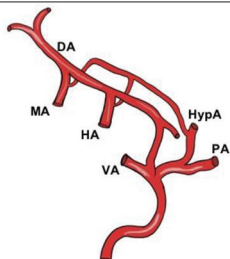
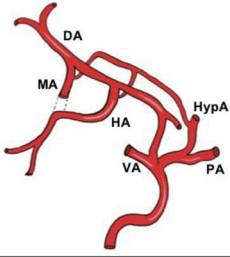
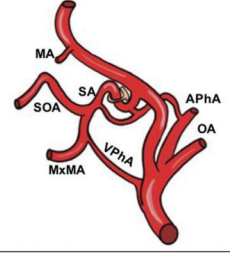
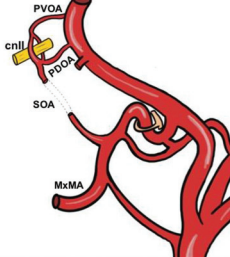
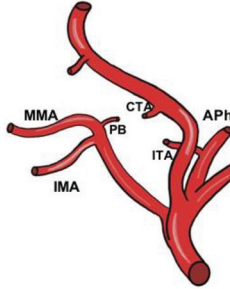
In the first stage of Padgett (4–5 mm embryos), the first 2 aortic arches initiate their natural regression, which allows the ICA to be individualized. The embryologic segments of the ICA are

derived from the third aortic arch and from the dorsal aorta cranial to the third aortic arch.^{31,32,39} The dorsal aorta also regresses at the same time between the third and fourth aortic arches. In terms of embryology, the ICA was divided by Lasjaunias and Santoyo-Vazquez⁵⁰ into 7 different segments. The first segment corresponds to the third aortic arch from the origin of the ventral pharyngeal artery (the future external carotid artery) to the junction between the third aortic arch and the dorsal aorta. The second segment is the dorsal aorta between the second and third aortic arch. The third segment is the dorsal aorta between the first and second aortic arch. The fourth segment is the dorsal aorta between the first aortic arch and the origin of the trigeminal artery (and the primitive maxillary artery). The fifth segment is the dorsal aorta between the origin of the trigeminal artery (and the primitive maxillary artery) and the origin of the primitive dorsal OA (the future inferolateral trunk). The sixth segment is between the origin of the primitive dorsal OA and the origin of the primitive ventral OA. The seventh segment is between the primitive ventral OA and the primitive carotid bifurcation.^{23,24,31,39}

It is important to note that the carotid bulb does not have the same embryologic origin as the other segments of the ICA. It originates from the pharyngo-occipital system, which easily explains variations in origin of the ascending pharyngeal and occipital arteries. Agenesis or abnormal regression of ≥ 1 segment of the ICA explains an intratympanic course of the ICA¹² (by pharyngo-carotid anastomosis) and also the different type of “reperfusion” in the case of ICA agenesis.³² The intratympanic course of the ICA, also named as an “aberrant flow of the ICA,” is the consequence of the abnormal regression of the first and second segments, with anastomosis between the inferior tympanic artery (from the ascending pharyngeal artery) and the caroticotympanic artery (from the carotid artery) that distally infuses the ICA. The pseudo-ICA, consequently, has an intratympanic course without passing through the stapes.^{12,39,51}

Hyostapedial System. “Hyostapedial artery” is the term used to describe the complete embryologic development of the second aortic arch. The SA, which develops from the hyoid artery and takes its name after passing through the crus of the stapes, is an embryonic artery present between Padgett³¹ stages III and VI (9–24 mm). This is an important embryologic system from which numerous dural, orbital, and facial arteries develop. The steps of SA development are summarized in Table 1.

Table 1: Major embryologic changes in the formation of the SA

Stage	Embryo size (mm)	Major evolutions	Graphic Illustration
I	4-5	- Regression of the second aortic arch - Hyoid artery (HA) formation (dorsal remnant of the 2nd aortic arch)	
II	5-6	- Elongation of the hyoid artery (HA) - Annexation of the mandibular artery (MA) territory (1st aortic arch) by the hyoid artery (2nd aortic arch)	
III/ IV	7-12 12-14	- Cranial growing of the hyoid artery (stapedial artery) passing into the middle ear (crus of the stapes) - Extension of the two branches of the stapedial artery: Supraorbital (SOA) and Maxillomandibular (MxMA)	
V	16-18	- Maximal development of the stapedial artery - Annexation of the supraorbital branch (SOA) by the ophthalmic artery (PDOA, PVOA) - Regression of the trans-osseous (superior orbital fissure) segment of the supraorbital branch	
VI	20-24	- Annexation of the maxillomandibular branch by the ventral pharyngeal artery - Regression of the intratympanic segment of the stapedial artery	

The hyoid artery is the dorsal remnant of the second aortic arch, which regresses early in the embryologic development (4-5 mm, Padgett³¹ stage I).^{2,12} After this arch regression, at approximately the 5–6-mm stage (Padgett³¹ stage II), the hyoid artery presents a rapid lateral elongation between the beginning of the 7th and 19th weeks of gestation, when it gives an anastomosis to the mandibular artery (remnant of the first aortic arch).³¹ During Padgett³¹ stage III (7–12-mm embryos), the hyoid artery is prominent and grows cranially, passing, as the SA, through the crus of the stapes. In contrast, the mandibular artery is short and difficult

to identify. During this stage, the initial forms of the 2 main branches of the SA (supraorbital and maxillomandibular divisions) are visible, even if they are definitively established only in Padgett³¹ stage IV (12–14 mm).

At the 16–18-mm stage (Padgett³¹ stage V), the SA continues its elongation to the gasserian region, passing into the future tympanic cavity and, particularly, through the crus of the stapes, reaching its maximum development.^{23,31,32} The supraorbital artery, which follows the ophthalmic root of the trigeminal nerve, allows the development of orbital branches (supraorbital,

Table 2: Summary of embryologic steps of the stapedia and ophthalmic systems

Stapedial System	Embryo Size (mm)	Ophthalmic System
Regression of the second aortic arch; hyoid artery formation (dorsal remnant of the second aortic arch)	4–5	Primitive maxillary artery as temporary branch; primitive dorsal OA appearance
Elongation of the hyoid artery; annexation of the mandibular artery (first aortic arch) by the hyoid artery (second aortic arch)	5–6	Primitive hyaloid artery = plexiform channels; primitive ventral OA appearance
Cranial growing of the hyoid artery (SA) passing into the middle ear (crus of the stapes)	7–14	Formation of primitive hyaloid and common ciliary arteries
Maximum development of the SA; extension of the 2 branches of the SA, supraorbital and maxillomandibular	16–19	Migration in origin of the primitive ventral OA; regression of the primitive dorsal OA; formation of the anastomotic ring
Regression of the transosseous (superior orbital fissure) segment of the supraorbital branch	20–24	Ventral interruption of the anastomotic ring; annexation of the supraorbital branch by the OA

lacrimal, ethmoids, and frontal arteries) and also of the MMA.³⁹ The maxillomandibular artery exits the cranial cavity through the foramen spinosum and, extracranially, gives its 2 branches: infra-orbital and mandibular arteries (the future infraorbital and inferior alveolar arteries). After this maximum development of the SA, 2 annexations and 2 regressions occur to give the adult configuration of the MMA. Intracranially, the orbital branches are annexed by the primitive OA and the trans-sphenoidal segment of the supraorbital branch regresses, leaving an anastomotic artery between the anterior branch of the MMA and the lacrimal artery (the sphenoidal artery), which penetrates the orbit through the superior orbital fissure. Extracranially, the ventral pharyngeal artery annexes the maxillomandibular artery of the SA, forming the proximal stem of the MMA and becoming the external carotid artery. Consequently, by flow reversal into the SA, its tympanic portion regresses and leaves, as remnants, the caroticotympanic artery (from the ICA) and the superior tympanic artery (from the petrous branch of the MMA). These annexations and regressions with regard to the SA happen during Padgett³¹ stage VI (20–24-mm embryos).^{31,39}

OA. The embryogenesis of the primitive OA is related in some phases to the hyostapedial systems. These phases are summarized and related to the concomitant SA steps of development presented in Table 2. The formation of the OA depends on 2 different arteries during the 4–18-mm stages; the primitive dorsal OA and the primitive ventral OA. The primitive dorsal OA develops from the cavernous segment of the primitive ICA and the primitive ventral OA develops from the anterior division of the primitive ICA. The primitive dorsal OA penetrates the orbit through the superior orbital fissure and the primitive ventral OA penetrates through the optic canal.

Next, 2 major anastomoses between these 2 arteries are formed. The first one is an intraorbital plexiform anastomosis supplied by the 2 arteries around the optic nerve (the future second segment of the OA). The second anastomosis is intradural between the primitive ventral OA and the primitive ICA to form the definitive supraclinoidal origin of the OA. The proximal portion of the primitive ventral OA (between its origin on the ACA and its anastomosis with the ICA), then regresses to give the adult stem of the OA.

In the following step, the proximal part of the primitive dorsal OA regresses and its remnant becomes the inferolateral trunk of the primitive ICA. Concomitantly, in 7- and 24-mm embryos (stages III–VI of Padgett), the SA arises from the primitive hyoid arch of the petrous ICA. As described above, its supraorbital artery enters the orbit to give orbital branches. It divides into 2 different branches: the ethmoidonasal and lacrimal arteries. The ethmoidonasal artery anastomoses with the primitive OA at the arterial ring around the optic nerve, as previously described. The trans-sphenoidal part of the supraorbital artery then regresses, and its orbital branches are annexed by the primitive OA to give the adult conformation.

Trigeminal Artery. The trigeminal artery represents one of the carotid-basilar anastomoses, which appears in 4–5-mm embryos (Padgett³¹ stage I) and disappears in 12-mm embryos (Padgett³¹ stage III). It originates from the basilar artery, between the superior and the antero-inferior cerebellar arteries, passes medial to the gasserian ganglion, and follows the trigeminal nerve to the primitive ICA at the level of the junction between the fourth and fifth segments of the ICA. The trigeminal artery has a common origin on the ICA with the primitive maxillary artery. The carotid remnant of these 2 primitive arteries will be the future meningo-hypophyseal trunk, from which the lateral clival, marginal tentorial, and inferior hypophyseal arteries will originate.^{31,32} The basilar remnant of the trigeminal artery could be considered responsible for MMA possible origin from the basilar artery. In fact, when this trigeminal remnant anastomoses with a persistent SA, the final MMA will arise from the BA. The details of this variant are explained in Part 2 of our article.

Formation of the Inferolateral Trunk. As supported by the theory of Lasjaunias,^{23,24,39} with regard to the embryology of the OA, the inferolateral trunk is the carotid remnant of the primitive dorsal OA. The primitive dorsal OA develops from the cavernous segment of the primitive ICA and penetrates the orbit through the superior orbital fissure. In 40-mm embryos, the proximal part of the primitive dorsal OA regresses, and its remnant becomes the inferolateral trunk. At the adult configuration, the inferolateral trunk is composed of 4 branches: 1) superior branch, which supplies the roof of the cavernous sinus; 2) anteromedial branch,

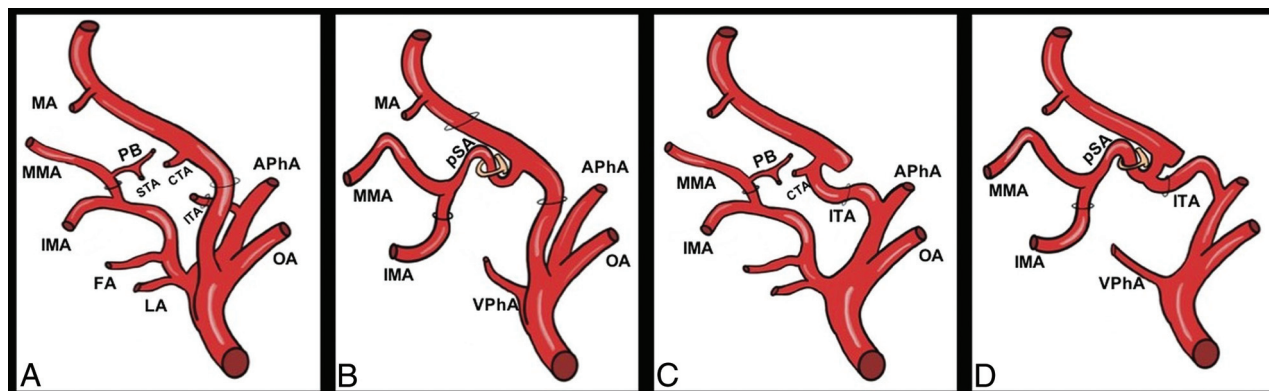


FIG 2. Persistent SA and intratympanic flow of the ICA. A, The normal regression process of the hyostapedial system. After the SA proximal regression, its proximal remnant originates from the ICA as the carotico-tympanic artery (CTA). Its distal part is annexed by the ventral pharyngeal artery, the future external carotid artery, to give the internal maxillary artery its first and major branch, the middle meningeal artery (MMA). The MMA side of the SA persists as the petrosal branch (PB), which gives birth to the superior tympanic artery (STA). The anastomotic branch to the hyostapedial system from the ascendant pharyngeal artery (APhA) persists as the inferior tympanic artery (ITA). The CTA, STA, and ITA contribute to the adult vascularization of the middle ear. B, Persistent SA. In the case of complete SA persistency, this artery originates from the petrous ICA and gives, intracranially, the MMA, and the internal maxillary artery exits the middle fossa skull base through the foramen spinosum as the extracranial branch. This variant is due to the lack of annexation of the maxillary-mandibular branch by the ventral pharyngeal artery. C, Pseudopetrous or aberrant intratympanic ICA. This variant is due to the agenesis of the cervical ICA (third aortic arch). The cervical ICA agenesis is bypassed through a hypertrophic inferior tympanic artery, which courses from the APhA through the inferior tympanic canal and reaches the CTA. The intratympanic ICA flow is called “aberrant” because the artery does not course into its usual canal but into an enlarged inferior tympanic canal. D, Pseudopetrous ICA with or without SA persistency. This variant is due to 2 phenomena: the agenesis of the cervical ICA and SA persistency. In this case, the ITA does not reach the CTA but the persistent SA (pSA).

which passes into the superior orbital fissure; 3) anterolateral branch, which runs into the foramen rotundum; and 4) posterior branch, which passes medial to the gasserian ganglion. These branches present numerous anastomoses in the cavernous region, which correspond to the remnants of the primitive trigeminal artery, OA, SA, and maxillary artery.¹⁵

Persistence of the SA

Changes in the events previously described can determine different adult configurations of the arteries related to the hyostapedial system, such as the persistence of the SA. The incidence of this variant is estimated to be 0.48% after a series of >1000 temporal bone dissections.⁶ The possible variants related to the persistence of the SA are illustrated in Fig 2, including the rarer association of SA persistence with the so-called pseudopetrous ICA.

Complete Persistence of the SA

The complete persistence of the SA is a very rare variant, and only 2 cases were published in the context of an ICA aneurysm²⁵ or PHACE (posterior fossa malformations, hemangioma, arterial anomalies, coarctation of the aorta/cardiac defects, and eye abnormalities) syndrome.³⁹ In these cases, the SA could be seen as in the embryo, taking its origin from the petrous ICA, passing through the middle ear, and giving its 2 branches: one intracranial, which corresponds to the MMA, and the other extracranial, leaving the cranial cavity through the foramen spinosum. Consequently, the foramen spinosum is enlarged, the cochlear promontory is eroded, and the internal maxillary artery arises from the SA instead of the external carotid artery. Such an anatomic variant could easily be explained by the embryology and, particularly, by the absence of annexation of the maxillomandibular branch (of the SA) by the ventral pharyngeal artery.

Consequently, in the absence of reversion of the arterial flow into the SA, its proximal (intratympanic) stem could not regress.³⁹

Partial Persistence of the SA

The partial persistence of the SA is more frequent^{1,7,21-24,26,32,52,53} and, in this case, only the intracranial branch of the SA keeps its origins from the petrous ICA. The foramen spinosum is absent (an orifice does not exist without its contents) or reduced in size and the MMA arises from the SA instead of the internal maxillary artery. This variant is explained by the regression of the proximal part of the maxillomandibular artery instead of the proximal part of the SA.³⁹ A rare case of a bilateral partial persistent SA is illustrated in Fig 3. We note that it is surprising that the complete persistence of the SA is much rarer than its partial persistence. The reasons of this phenomenon are still unknown, and are probably related to some unknown phases of the annexation of the maxillomandibular branch by the ventral pharyngeal artery.

Persistence of the SA Associated with Aberrant Flow of the ICA (Pseudo-Petrous ICA)

In rare cases,^{12,21,23,24,54,55} the persistence of the SA is associated with an intratympanic course of the ICA (also known as “aberrant flow of the ICA”). This association of vascular variants with regard to both carotid and hyostapedial systems was described anatomically and angiographically.^{32,39} In these cases, the ICA enters the skull base through an enlarged inferior tympanic canal (narrowing of the vessel on DSA), passes into the tympanic cavity to bend anteriorly, and reaches its normal carotid canal. The MMA arises from the ICA in its tympanic segment and passes through the stapes to have the same course described in the previous paragraph. The exocranial orifice of the carotid canal,

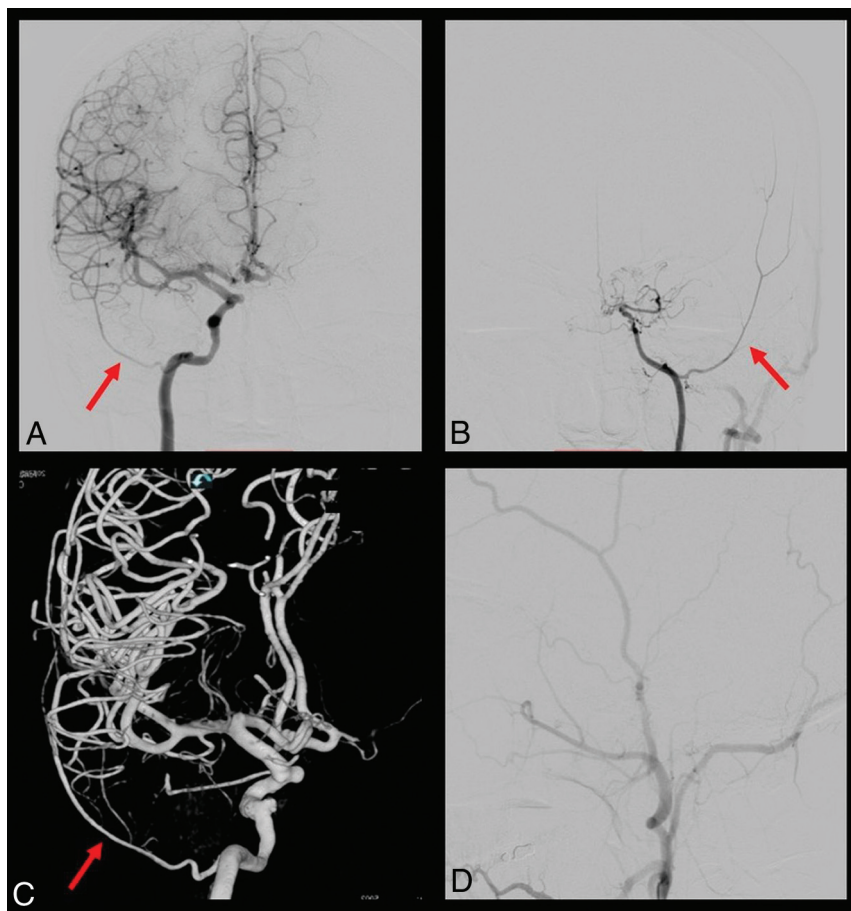


FIG 3. DSA with a bilateral partial persistent SA in a patient with Moyamoya disease. A and B, show the right and left ICA injection, respectively, with the MMA (red arrow) arising from the petrous ICA. This phenomenon is possible because of bilateral SA persistency. B, The ICA is narrowed and stopped at the level of the OA due to Moyamoya disease. C, A 3D reconstruction of the right ICA from which the MMA originates (red arrow). D, The left external carotid artery injection without the MMA enhancement because of its absence.

therefore, is absent in these cases.^{23,24} The intratympanic course of the ICA is explained by the agenesis of the first 2 segments of the primitive ICA. The cervical segment, in fact, is the ascending pharyngeal artery with a hypertrophied inferior tympanic artery that maintains its anastomosis with the carotico-tympanic artery (the branch of the ICA) into the tympanic cavity. The correct term for this aberrant flow of the ICA, in reality, is a tympano-carotico-tympanic variant.³⁹ It bypasses the absence of the cervical ICA and provides a collateral circulation to the remainder of the ICA. Therefore, the cervical and intratympanic segments of this artery do not derive from the carotid system but from the pharyngo-occipital and hyostapedial systems (pseudo-ICA).^{23,24,39} The agenesis of the first 2 segments of the ICA could be partial, and the ICA seems duplicated.⁵⁴

Pharyngo-Tympano-SA

This very rare variant was first described by Lasjaunias et al,²³ in 1977, in their original publication. The same case served as an illustration in the textbook, *Surgical Neuroangiography*,³⁹

and only 1 similar case was published by Baltsavias et al.⁴² The MMA arises from the cervical portion of the ICA, ascends along the cervical ICA, enters into the tympanic cavity through the inferior tympanic canal, and follows the usual course of the SA. The 2 cases described were presented as “partial” persistence of the SA, with only the MMA arising from the SA, and the absence of the foramen spinosum. In this variant, an annexation of the SA by the inferior tympanic artery (the branch of the ascending pharyngeal artery), with regression of the proximal part of the SA explains this vascular configuration. Therefore, the SA arises from the cervical instead of the petrous segment of the ICA.

MMA Origin of the OA

The first description of orbital branches arising from the MMA was published in 1872 by Curnow.⁵⁶ He already noted in his case dissection that all orbital branches were supplied by the MMA except the central retinal artery, which kept its origin from the supracavernous ICA. Twenty years after this first description, Musgrove⁵⁷ presented another case of an MMA origin of the OA, also where the central retinal artery was supplied by the MMA and the ICA does not give any orbital artery, thus the orbit is vascularized by the MMA. After other isolated cases found during cerebral or orbital dissections,^{1,27,58,59}

Hayreh and Dass⁶⁰ reported 6 cases among their 170 orbital dissections in which the MMA participated completely (2 cases) or partially (4 cases) in the orbital supply. Moret et al³⁶ and Lasjaunias et al²³ described with precision the contribution of the MMA to the orbital vascularization and gave us a more comprehensive explanation of this anatomic variation.

The incidence of OA that arises from the MMA and penetrates the orbit through the superior orbital fissure is estimated as between 1.4 and 2.5%.⁶⁰ In this anatomic variation, orbital arteries are supplied by the anterior division of the MMA passing through the superior orbital fissure or through the spheno-orbital foramen. The central retinal artery generally keeps its vascular supply from the supracavernous ICA, but, in a few cases, the central retinal artery is also supplied by the MMA without ICA participation.⁶¹ A case of a complete supply of the orbital arteries by the MMA is illustrated in Fig 4.

As explained before, during the embryogenesis, the SA gives the supraorbital branch, which allows the development of the lacrimal artery and anastomoses with the OA around the optic nerve

(participation of the perioptic circle). The proximal part of the supraorbital branch then normally regresses and the lacrimal artery is annexed by the OA.³⁹ The persistence of MMA branches (or, in extremis, the MMA origin of the OA) could be explained by 2 different phenomena: the absence of an anastomosis between the supraorbital branch and the OA, with a consequent persistence of a dual supply of the orbital arteries, or the persistence of the proximal stem of the supraorbital branch of the SA, with regression of the primitive OA (complete MMA origin of the OA). Although the supraorbital branch bifurcates (lacrimal artery laterally and ethmoido-nasal medially) usually inside the orbit, in 30% of cases, it can divide outside the orbit in the middle cranial fossa. In such cases, the ethmoido-nasal artery passes through the superior orbital fissure, but

the lacrimal artery penetrates the orbit through its own canal, the speno-orbital foramen (canal of Hyrtl, lacrimal foramen, sinus canal). The medial branch, passing through the superior orbital fissure, is named the recurrent meningeal artery in its intraorbital segment and sphenoidal artery in its intracranial segment.^{23,31,39,60}

CONCLUSIONS

All the anatomic variations linked to the development of the SA, summarized in Table 3, could have a clinical impact. Because each vascular segment that regresses during embryologic life usually persists as an anastomosis between 2 arteries, the knowledge of the embryology and the presence of these anastomoses is of paramount importance in case of external carotid branch embolization. Each residual anastomosis represents a potential undesired leak of liquid embolic agent, with possible clinical implications. Knowledge of the embryologic variants described in this article represents the basis to understand the different possible adult configurations of the MMA and their clinical implications. The most common pathologies, whose treatment requires a detailed knowledge of MMA embryology and variants, are dural AVFs, meningeal tumors, and chronic subdural hematomas. Our purpose, therefore, is to give a detailed description of these variants and their clinical implications in the second part of our article.

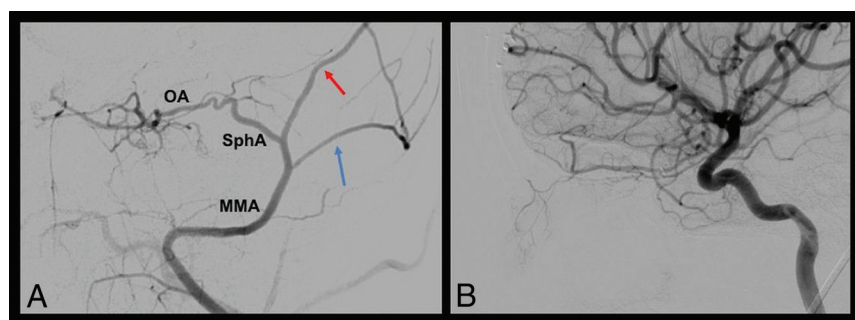


FIG 4. The MMA origin of the OA: a case of a complete OA origin from the MMA. The selective injection of the MMA in the anteroposterior projection (A) shows that the sphenoidal artery (SphA) arises from the MMA after its bifurcation into the anterior (red arrow) and posterior division (blue arrow). It courses along the inner surface of the temporal and sphenoidal bone and enters the orbit through the superior orbital fissure (SOF) to give rise to the OA. B, An oblique ICA injection, highlighting the absence of the OA.

Table 3: Anatomic variations implicating the SA

Anatomic Variations			Embryologic Implications	
Type	Incidence	Direct and Indirect Signs	Embryologic Explanation	Embryo Size (mm)
Complete persistence of the SA	2 cases	Petrous ICA origin of the MMA; petrous ICA origin of the internal maxillary artery; enlarged foramen spinosum; eroded cochlear promontory	Lack of annexation of the maxillomandibular branch by the ventral pharyngeal artery; persistence of the tympanic segment of the SA	24
Partial persistence of the SA	0.4%	Petrous ICA origin of the MMA; absent foramen spinosum; enlarged facial canal	Regression of the proximal segment of the maxillomandibular branch; persistence of the tympanic segment of the SA	24
Tympano-carotico-tympanic variant; aberrant flow of the ICA plus persistent SA	5 cases	Intratympanic course of the ICA; petrous ICA origin of the MMA; enlarged inferior tympanic canal; absent exocranial carotid canal	Regression of the proximal segment of the maxillomandibular branch; persistence of the tympanic segment of the SA; agenesis of the first 2 segments of the ICA; anastomosis between inferior tympanic artery and caroticotympanic artery	24; 4–5
Pharyngo-tympano-stapedial variant	2 cases	Cervical ICA origin of the MMA; absent foramen spinosum	Regression of the proximal segment of the maxillomandibular branch; persistence of the tympanic segment of the SA; anastomosis between the inferior tympanic artery and the caroticotympanic artery	24
MMA origin of the OA	n = 2%	MMA origin of the OA; no OA from the ICA or only the central retinal artery	Lack of annexation of the supraorbital branch by the OA; persistence of the sphenoidal part of the supraorbital branch	24

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